

[Reprinted from the Journal of the American Chemical Society, 74, 5487 (1952).]

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Physical Constants and Infrared Spectra of a Homologous Series of Dialkyl Sebacates

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The 18 diesters of sebacic acid with normal alcohols from methyl to octadecyl were prepared. Melting points, densities and refractive indices are given and the characteristics of the infrared absorption spectra are briefly discussed.

The compounds representing the homologous series of di-n-alkyl sebacates (dialkyl decanedioates) were synthesized primarily for use as substrates for the growth of certain bacteria and fungi as part of a study of the microbiological degradation of plasticizers. Ayailable data on these esters are scant; with the exception of di-n-octyl sebacate, no physical constants for the esters above din-propyl sebacate appear in the literature. Although the first three members of the series have been described before, they are included here for the sake of completeness.

Experimental

Tridecyl, pentadecyl, hexadecyl, heptadecyl and octadecyl alcohols³ were generously donated by Dr. E. Emmett Reid of Johns Hopkins University; nonyl alcohol was furnished by Dodge and Olcott, New York, with a specification of 98% purity, n^{20} 1.4340; tetradecanol was donated by Armour and Co., with a specification of 99% purity, m.p. 37.7°; hexyl alcohol was redistilled Eastman Kodak Co. technical grade, using the fraction boiling between 155-156°; all other alcohols were Eastman "white label" grade; methyl, ethyl and propyl were redistilled and all others used without further purification.

The sebacates were prepared by esterification of sebacic acid (Eastman "white label," recrystallized three times, m.p. 133.5°), with the appropriate alcohol in the presence of p-toluenesulfonic acid as catalyst. The crude product was dissolved in ether and thoroughly washed with 10% sodium bicarbonate. Those esters up to di-n-nonyl inclusive were distilled in vacuo, while the higher ones were further purified by repeated crystallization from acetone until

successive crops showed constant melting point. A further purification of all products was achieved by chromatographing them as ether solutions through a column packed with freshly activated adsorbents. For adsorption of acidic impurities, anhydrous alumina was used; for alcohols, siliegel; and for color and odor, fuller's earth. The liquid esters were finally subjected to a high vacuum for several hours to remove any remaining traces of solvent. All resulting products showed an acid number < 0.05.

Melting points of the esters up to di-n-nonyl inclusive were taken by freezing in a test-tube containing a narrow range thermometer, stirring constantly upon partial melting, and noting the melting point at the disappearance of the last crystal. Melting points of the esters above di-n-nonyl were taken by means of a calibrated Kosler micro hot-stage melting-point apparatus.

Results and Discussion

The esters up to and including di-n-nonyl sebacate at room temperature were colorless liquids of relatively low viscosity and with boiling points above 290°. The higher esters were white waxy crystalline solids.

Melting points and carbon-hydrogen analyses for the eighteen sebacates are listed in Table I. It is pertinent to point out that if these melting points are plotted as a function of carbon atoms in the alkyl radical, the curve is comparable to one obtained for the corresponding adipates (hexanedioates) by Feagan and Copenhaver, which exhibited a definite alternation in melting point values between odd and even alkyls. In the present case the same general phenomenon is noticeable, particularly above C₁₁. A curve of the values for odd-

(4) R. A. Feagan and J. E. Copenhaver, ibid., 62, 869 (1940).

⁽¹⁾ W. H. Stahl and H. Pessen, Applied Microbiology, in press.

⁽²⁾ J. T. Davies, Trans. Faraday Soc., 44, 909 (1948).

⁽³⁾ J. D. Meyer and E. E. Reid, This Journal, 55, 1574 (1933).

TABLE I

MELTING POINTS AND ANALYSES OF THE DIALKYL SEBA-

		CALES			
Diester	M.p. (cor.), °C.	Carbo Theor,	on, % Found	Hydro Theor.	gen, % Found
Methyl	23.8	62.65		9.64	
Ethyl	+1.5	65.08	•	10.15	
Propyl	0.0	67.09		10.56	
Butyl	-9.2	68.75	68.84	10.90	11.22
Pentyl	-3.0	70.18	69.86	11.19	11.18
Hexyl	+1.2	71.30	71.17	11.42	11.60
Heptyl	18.4	72.31	72.29	11.63	11.56
Octyl	18.6	73.19	73.06	11.81	11.86
Nonyl	26.2	73.96	74.08	11.97	11,91
Decyl	31.5	74.63	74.74	12.11	12.40
Undecyl	45.2	75.24	75.50	12.23	12.31
Dodecyl	49.1	75.78	75.70	12.34	12.55
Tridecyl	53.2	76.27	76.50	12.44	12.34
Tetradecyl	53.5	76,71	76.97	12.53	12.75
Pentadecyl	59.1	77.11	77.43	12.62	12.89
Hexadecyl	60.0	77.48	77.42	12.69	12.71
Heptadecyl	65.0	77.81	77.89	12.76	12.93
Octadecyl	66.3	78.12	78.21	12.82	12.97

carbon alkyls is slightly above that for even ones; this behavior is analogous to that of esters of monocarboxylic acids but is opposite to that of alcohols and free monocarboxylic and dicarboxylic acids.6

Densities, refractive indices and molar refractions for those esters which are liquid near room temperature (i.e., up to nonyl) are listed in Table II.

Infrared spectrograms for the eighteen esters were obtained; those for C₁, C₅ and C₁₈ alkyl esters are reproduced in Fig. 1. Except for some differences among the initial members of the series, the spectra appear conspicuously similar, all having absorption bands of nearly equal intensity at the same wave

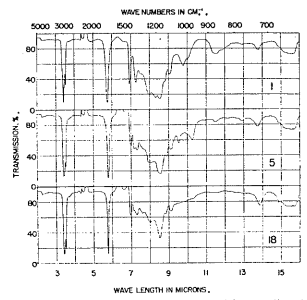


Fig. 1.—Infrared spectra of dimethyl, di-(n-pentyl) and di-(n-octadecyl) sebacates. Baird spectrograph with NaCl prism; 10.10% (C1), 10.34% (Cs), 8.80% (C18) in carbon disulfide solution; cell thickness, 0.1 mm.

TABLE II

DENSITY	AND	REFRACTIVE	INDEX	OF	CERTAIN	DIALKYL	
Sebacates							

			,			
Diester		des,a	n ²⁵ D (Abbe) <i>b</i>	Mol, refr. c Theor. Found		
	Methyl	0.9883	1.4368	60.92	61.02	
	Ethyl	.9590	1.4348	70.15	70.27	
	Propyl	.9451	1.4374	79.39	79.45	
	Butyl	,9321	1.4397	88.63	88.91	
	Pentyl	.9266	1.4426	97.86	97.92	
	Hexyl	.9183	1.4445	107.10	107.29	
	Heptyl	.9126	1,4460	116.33	116.48	
	Octyl	.9074	1.4480	125.57	125.87	
	Nonyl	.9040	1.4492	134.81	134.92	

 a Δd^{25} , per degree = -0.0004. b Δn D per degree = -0.0004. "Physical Methods of Organic Chemistry," A. Weissberger, ed., Interscience Publishers, Inc., New York, N. Y., 1945, p. 673.

lengths. Among the most outstanding of these are the following, with indicated assignments based on published frequency charts⁷⁻¹⁰: a strong, narrow doublet at 3.44 and 3.52 μ (methylene stretching), another strong, narrow band at 5.77 (aliphatic ester carbonyl stretching), bands at 6.94 and 7.31 μ (methylene and methyl bending), a group of bands or inflections at 7.60, 8.00, 8.50, $8.\overline{90}$ and 9.12μ (methylene twisting; ether linkage), and a weaker but distinctive band at 13.82 μ (apparently indicative of continuous chains of four or more carbon atoms due to methylene wagging11,12 observed in particular for dicarboxylic acids above

It is of interest to note that when the total number of alkyl carbons begins to exceed the number in the acid chain (i.e., beyond the dipentyl ester) the bands in the 9 to 12 μ region begin to become similar and remain so for all the higher homologs. The reason for this may be that the shorter alkyl chains are subject to special coupling effects which at first differ in character from one member of the series to the next, as evidenced by markedly different additional vibrations. Once the alkyl chains have reached a certain length these effects might be expected to remain constant and any further changes in the spectrum to be due solely to the effect of subsequent methylene groups attached to the chain at a point relatively far removed from the ester linkage.

By way of comparison with esters of monocarboxylic acids it may be pointed out that, although the molecules here dealt with are all symmetric, no special symmetry effects7 should be expected since the acid carbon chain separating the two alkyl portions of the molecule is so long (C_{10}) that the alkyl chains vibrate virtually independently of each other.

The homologous series exhibits a small but defi-

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 (11) C. Schaefer and F. Matossi, "Das Ultrarote Spektrum," Julius Springer, Berlin, 1930, p. 268.
- (12) O. D. Shreve, M. R. Heether, H. B. Knight and Daniel Swern, Anal. Chem., 22, 1948 (1950).
 - (13) J. Lecomte, Compt. rend., 211, 778 (1940).

⁽⁵⁾ J. H. Hoback, et al., This Journal, 65, 1606 (1943).

⁽⁶⁾ L. F. Fieser and M. Fieser, "Organic Chemistry," D. C. Heath and Co., Boston, Mass., 1950, pp. 302, 409.

nite progression in intensities of certain bands. As the alkyl chain length increases from 1 to 18 carbons, the intensity of the entire group of bands from 6.94 to 9.12 μ decreases from, e.g., 46% transmission at 9.12 μ to 68%. This suggests a possible aid in the identification of the alkyl radical of an unknown sebacate by comparison of its % transmission under specified conditions at selected wave

lengths with values taken from prepared plots of % transmission vs. number of skeletal carbon atoms at corresponding wave lengths.

Acknowledgment.—The authors are grateful to Miss D. A. Wiener for obtaining the infrared spectrograms and Mr. Sam Cohen for his help in their interpretation.

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